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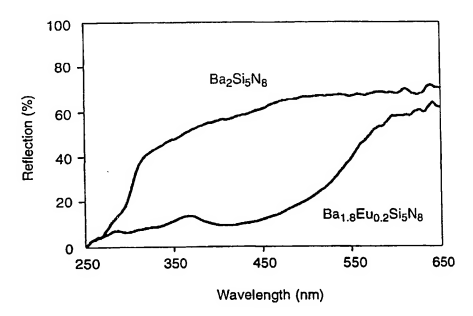
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(54) Title: PIGMENT WITH DAY-LIGHT FLUORESCENCE



WO 01/39574 A1 (57) Abstract: A pigment, especially a yellow to red emitting luminescent material, with a host lattice of the nitridosilicate type $M_x Si_y N_z$: Eu, wherein M is at least one of an alkaline earth metal chosen from the group Ca, Sr, Ba and wherein z = 2/3x + 4/3y.

Pigment with day-light fluorescence

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Technical Field

This invention relates to a Pigment with day-light fluorescence and more particularly, but not exclusively to a pigment absorbing blue to green light and emitting fluorescence within the yellow to red spectral region under excitation by daylight or by an artificial light source. Further absorption in other spectral regions is possible, especially in the UV. More specifically, such a pigment can be used as a phosphor for light sources, especially for Light Emitting Diodes (LED) or electrical lamps. The pigment belongs to the class of rareearth activated silicon nitrides.

Background Art

For Eu²⁺ -doped material normally UV-blue emission is observed (Blasse and Grabmeier: Luminescent Materials, Springer Verlag, Heidelberg, 1994). Several studies show that also emission in the green and yellow part of the visible spectrum is possible (Blasse: Special Cases of divalent lanthanide emission, Eur. J. Solid State Inorg. Chem. 33 (1996), p. 175; Poort, Blokpoel and Blasse: Luminescence of Eu²⁺ in Barium and Strontium Aluminate and Gallate, Chem. Mater. 7 (1995), p. 1547; Poort, Reijnhoudt, van der Kuip, and Blasse: Luminescence of Eu²⁺ in Silicate host lattices with Alkaline earth ions in a row, J. Alloys and Comp. 241 (1996), p. 75). Hitherto, red Eu²⁺ luminescence is observed only in some exceptional cases, such as in alkaline earth sulphides and related lattices of the rock-salt type (Nakao, Luminescence centers of MgS, CaS and CaSe Phosphors Activated with Eu²⁺ Ion, J. Phys. Soc. Jpn. 48(1980), p. 534), in alkaline earth thiogallates (Davolos, Garcia, Fouassier, and Hagenmuller, Luminescence of Eu²⁺ in Strontium and

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Barium Thiogallates, J. Solid. State Chem. 83 (1989), p. 316) and in some borates (Diaz and Keszler; Red, Green, and Blue Eu²⁺ luminescence in solid state Borates: a structure-property relationship, Mater. Res. Bull. 31 (1996), p. 147). Eu²⁺ luminescence in alkaline-earth silicon nitrides has hitherto only been reported for MgSiN₂:Eu (Gaido, Dubrovskii, and Zykov: Photoluminescence of MgSiN₂ Activated by Europium, Izv. Akad. Nauk SSSR, Neorg. Mater. 10 (1974), p. 564; Dubrovskii, Zykov and Chernovets: Luminescence of rare earth Activated MgSiN₂, Izv. Akad. Nauk SSSR, Neorg. Mater. 17 (1981), p. 1421) and Mg_{1-x}Zn_xSiN₂:Eu (Lim, Lee, Chang: Photoluminescence Characterization of Mg_{1-x}Zn_xSiN₂:Tb for Thin Film Electroluminescent Devices Application, Inorganic and Organic Electroluminescence, Berlin, Wissenschaft und Technik Verlag, (1996), p. 363). For both Eu²⁺ luminescence in the green and green/blue part of the spectrum was found.

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New host lattices of the nitridosilicate type are based on a three dimensional network of cross-linked SiN₄ tetrahedra in which alkaline earth ions (M= Ca, Sr and Ba) are incorporated. Such lattices are for example Ca₂Si₅N₈ (Schlieper and Schlick: Nitridosilicate I, Hochtemperatursynthese und Kristallstruktur von Ca₂Si₅N₈, Z. anorg. allg. Chem. 621, (1995), p. 1037), Sr₂Si₅N₈ and Ba₂Si₅N₈ (Schlieper, Millus and Schlick: Nitridosilicate II, Hochtemperatursynthesen und Kristallstrukturen von Sr₂Si₅N₈ and Ba₂Si₅N₈, Z. anorg. allg. Chem. 621, (1995), p. 1380), and BaSi₇N₁₀ (Huppertz and Schnick: Edge-Sharing SiN₄ tetrahedra in the highly condensed Nitridosilicate BaSi₇N₁₀, Chem. Eur. J. 3 (1997), p. 249). The lattice types are mentioned in Table 1.

Sulfide based phosphors (e.g. earth alkaline sulfides) are less desirable for lighting applications, especially for LED applications, because they interact with the encapsulating resin system, and partially suffer from hydrolytic attack. Red emitting Eu²⁺ activated borates show already temperature quenching to a certain degree at the operating temperature of LEDs.

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Disclosure of the Invention

It is, therefore, an object of this invention to obviate the disadvantages of the prior art. It is another object of the invention to provide a pigment for day-light fluorescence. It is a further abject to provide a yellow to red emitting luminescent material which is excitable at wavelengths around 200 to 500 nm, preferably 300 to 500 nm, together with high chemical and thermal stability.

Especially high stability up to at least 100 °C is highly desirable for LED applications. Their typical operation temperature is around 80 °C.

These objects are accomplished by the characterising features of claim 1. Advantageous embodiments can be found in the dependant claims.

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The new pigments show at least absorption within the blue-green spectral region. Furthermore they show fluorescent emission under absorption. Those Eu²⁺-doped luminescent materials show emission within the yellow to red spectral region, especially long wavelength red, orange or yellow emission. These pigments are based on alkaline-earth silicon nitride material as host-lattices. They are very promising, especially for LED applications, when used as phosphors. Hitherto white LEDs were realised by combining a blue emitting diode with a yellow emitting phosphor. Such a combination has only a poor colour rendition. A far better performance can be achieved by using a multicolour (for example red-green-blue) system. Typically the new material can be used together with a green-emitting (or yellow-emitting) phosphor, for example strontiumaluminate SrAl₂O₄:Eu²⁺, whose emission maximum is around 520 nm.

In detail, the new Pigment with day-light fluorescence uses a host lattice of the nitridosilicate type $M_xSi_yN_z$:Eu, wherein M is at least one of an alkaline earth metal chosen from the group Ca, Sr, Ba and wherein z = 2/3x + 4/3y. The incorporation of nitrogen increases the proportion of covalent bond and ligand-field splitting. As a consequence this leads to a pronounced shift of

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lattices.

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excitation and emission bands to longer wavelengths in comparison to oxide

Preferably, the pigment is of the type, wherein x = 2, and y = 5. In another preferred embodiment, the pigment is of the type, wherein x = 1, and y = 7.

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- Preferably, the metal M in the pigment is strontium because the resulting phosphor is emitting at relatively short yellow to red wavelengths. Thus the efficiency is rather high in comparison to most of the other elected metals M.
 - In a further embodiment the pigment uses a mixture of different metals, for example Ca (10 atom.-%) together with Ba (balance), as component M.
- These materials show high absorption and good excitation in the UV and blue visible spectrum (up to more than 450 nm), high quantum efficiency and low temperature quenching up to 100 °C.

It can be used as a pigment for coloring goods or as a phosphor for luminescence conversion LEDs, especially with a blue light emitting primary source together with one or more other phosphors (red and green).

Brief Description of the Drawings

- Fig. 1: Diffuse reflection spectra of undoped Ba₂Si₅N₈ and Ba₂Si₅N₈:Eu;
- Fig. 2: Diffuse reflection spectra of undoped BaSi₇N₁₀ and BaSi₇N₁₀:Eu;
- Fig. 3: Emission spectrum of Ba₂Si₅N₈:Eu;
- Fig. 4: Emission spectrum of BaSi₇N₁₀:Eu;
- 20 Fig. 5-7: Emission spectrum of several embodiments of Sr₂Si₅N₈:Eu;
 - Fig. 8: Emission spectrum of Ca₂Si₅N₈:Eu.

Detailed Embodiments

 Eu_2O_3 (with purity 99,99 %), or Eu metal (99,99 %), Ba metal (> 99 %); Sr metal (99 %), Ca_3N_2 (98 %), or Ca powder (99,5%) and Si_3N_4 (99,9 %) were used as commercially available starting materials. Ba and Sr were nitrided by

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firing at 550 and 800 °C under a nitrogen atmosphere. Subsequently, Ca_3N_2 or nitrided Ba, Ca or Sr were ground in a mortar and stoichiometrically mixed with Si_3N_4 under nitrogen atmosphere. The Eu-concentration was 10 atom.-% compared to the alkaline earth ion. The powdered mixture was fired in molybdenum crucibles at about 1300-1400 °C in a horizontal tube furnace under nitrogen/hydrogen atmosphere. After firing, the powders were characterised by powder X-ray diffraction (Cu, K α -line), which showed that all compounds had formed.

The undoped $Ba_2Si_5N_8$, $Ca_2Si_5N_8$ and $BaSi_7N_{10}$ are greyish-white powders. These undoped rare-earth activated silicon nitrides show high reflection in the visible range (400-650 nm) and a strong drop in the reflection between 250-300 nm (Fig. 1 and 2). The drop in reflectance is ascribed to host-lattice absorption. The Eu-doped samples are orange-red, except for $BaSi_7N_{10}$:Eu which is orange-yellow (Table 1). The strong coloration is unique for Eu^{2+} -doped rare-earth activated silicon nitrides and make these material interesting orange-red pigments. A typical example of a reflection spectrum of $Ba_2Si_5N_8$:Eu shows that the absorption due to Eu is superposed on the host-lattice absorption and extends up to 500-550 nm (Fig. 1). This explains the red-orange colour of these compounds. Similar reflection spectra were observed for $Sr_2Si_5N_8$:Eu and $Ca_2Si_5N_8$:Eu.

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For $BaSi_7N_{10}$:Eu the absorption of Eu is less far in the visible part (Fig. 2), which explains the orange-yellow colour of this compound.

All samples show efficient luminescence under UV excitation with emission maxima in the orange-red part of the visible spectrum (see Table 1). Two typical examples of emission spectra can be seen in Figs. 3 and 4. They show that the emission is at extremely long wavelengths (for Eu²⁺ emission) with maxima up to 660 nm for BaSi₇N₁₀:Eu (Fig. 4.). Excitation bands are observed at low energy which is the result of a centre of gravity of the Eu²⁺ 5d band at low energy and a strong ligand-field splitting of the Eu²⁺ 5d band, as can be expected for N³⁻ containing lattices (van Krevel, Hintzen, Metselaar,

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and Meijerink: Long Wavelength Ce³⁺-luminescence in Y-Si-O-N Materials, J. Alloys and Comp. 168 (1998) 272).

Since these materials can convert blue into red light due to low-energy excitation bands, they can be applied in white light sources, for example based on primarily blue-emitting LED's (typically GaN or InGaN) combined with red, yellow and/or green emitting phosphors.

Table 1:

Compound	Crystal structure	Color	Emission Maximum (nm)*
Ca ₂ Si ₅ N ₈ :Eu	Monoclinic	Orange-Red	600 to 630
Sr ₂ Si ₅ N ₈ :Eu	Orthorhombic	Orange-Red	610 to 650
Ba ₂ Si ₅ N ₈ :Eu	Orthorhombic	Orange-Red	620 to 660
BaSi ₇ N ₁₀ :Eu	Monoclinic	Orange-Yellow	640 to 680

^{*}depending on the conditions for preparation and concentration of the activator; typical values for Eu-concentration may vary between 1 and 10% compared to the alkaline-earth ion M

These emission maxima are unusually far in the long wavelength side. A specific example is a phosphor of the type Sr_{1.8}Eu_{0.2}Si₅N₈. Its emission spectrum is shown in fig. 5.

Another embodiment for realising M is the use of Zn. It can replace Ba, Sr or Ca fully or partially.

A further embodiment for replacing Si fully or partially is Ge. An concrete embodiment is Sr_{1.8}Eu_{0.2}Ge₅N₈.

Some further specific examples were investigated:

The preparation conditions and optical properties of the red emitting phosphor $Sr_2Si_5N_8$: Eu^{2+} were investigated. Optimisation showed a quantum efficiency of about 70 %. The emission is tuneable between 610 and 650 nm, depending on the Eu^{2+} concentration in the sample and the heating conditions. The absorption at 400 nm and 460 nm is high (reflection of only 15-40 %) and the temperature quenching of the luminescence at 80° C is low (only 4 %). The particle size of the phosphor is

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without milling below 5 µm. These properties make this phosphor very interesting especially for application in both the UV and blue LED.

For the nitride synthesis, the starting materials are Si₃N₄ (99,9% (mainly α-phase), Alfa Aesar), Sr metal (dendritic pieces 99,9 %, Alfa Aesar) and Eu₂O₃ (4N). The Sr metal has to be nitrided and in case one uses instead of Eu₂O₃ Eu metal, this has also to be nitrided.

The Sr metal is milled by hand in an agath mortar in an argon glovebox and nitrided at 800°C under N₂. This results in a nitration over 80 %.

After remilling, the nitrided metal, together with Si₃N₄ and Eu₂O₃, is milled and mixed by hand again in the glovebox. The heating of this mixture has typically the following parameters:

18°C/min to 800°C
5h at 800°C
18°C/min to T_{end} (1300-1575°C)
5h at T_{end} (1300-1575°C)
H₂(3.75%)/N₂ 400l/h

Ca₂Si₅N₈:Eu²⁺ samples were made with Ca₃N₂ as starting material.

An overview of all the samples is given in table 1. Typically, the samples were first heated at 800°C, and then they were heated a second time in the same cycle at elevated (1300-1600°C) temperatures. The samples were then milled (mill under air), sieved and measured.

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Table 1: parameters of heating cycles of (Ca,Sr)₂Si₅N₈:Eu²+ samples

Code	Ca/sr	Eu ²⁺ (%)	Time 1 (h)	Temp. 1 (°C)	Time 2 (h)	Temp. 2 (°C)
EC/HU 31/00	Ca	10	5	800	5	1400
EC/HU 42/00	Ca	1	5	800	5	1565
EC/HU 41/00	Ca0.4Sr1.4	10	5	800	5	1565
EC/HU 62/00	Sr	1	5	800	5	1400
EC/HU 63/00	Sr	2	5	800	5	1400
EC/HU 64/00	Sr	3	5	800	5	1400
EC/HU 65/00	Sr	5	5	800	5	1400
EC/HU 66/00	Sr	8	5	800	5	1400
EC/HU 67/00	Sr	10	5	800	5	1400

The samples that are obtained after this heating show a color of deep orange for $10\%~Eu^{2+}$ containing $Sr_2Si_5N_8$ samples. With less Eu^{2+} the colour is fainter. The Ca samples have a yellow-orange colour.

There is also another interesting feature: the powder particles are very small with an average particle size d_{50} between about 0,5 and 5 μ m, a typically value is d_{50} =1.3 μ m. The small particle sizes are advantageous for the processing of LEDs with luminescent material. For example they allow a homogeneous distribution in the resin.

10 Table 2: Optical data of (Ca,Sr)₂Si₅N₈:Eu²+ samples

Code	Ca/Sr	Eu ²⁺ (%)	Em. Max (nm)	Refl. 400 (%)	Refl. 460 (%)	QE (%)	х	у
EC/HU 31/00	Ca	10	619	12	19	26	0.600	0.396
EC/HU 42/00	Ca	1	603	47	58	37	0.555	0.435
EC/HU 41/00	Ca0.4 Sr1.4	10	660	17	22	59	0.636	0.,354
EC/HU 62/00	Sr	1	609	53	58	70	0.602	0.393
EC/Hu 63/00	Sr	2	618	43	48	73	0.615	0.381
EC/Hu 64/00	Sr	3	621	36	41	72	0.622	0.374
EC/Hu 65/00	Sr	5	624	26	32	67	0.632	0.365
EC/HU 66/00	Sr	8	636	21	26	67	0.641	0.356
EC/HU 67/00	Sr	10	644	17	22	64	0.642	0.354

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Concerning table 2 all samples were typically first heated in a first cycle (for example 800°C for 5h), as already outlined above.

Included in table 2 are the position of the emission maximum, the mean wavelength, the reflection at 400 and 460 nm, the quantum efficiency and finally the x and y colour coordinates.

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From table 2 it can be derived that the pure Ca samples are not as favourable as the Sr samples. It is surprising that the Sr-Ca compound has an emission wavelength that is larger than that of the pure Sr compound.

Specific examples are shown in Figures 6 to 8. Figure 6 shows the energy distribution (in arbitrary units) and reflection (in percent) of sample HU 64/00 (Sr₂Si₅N₈:Eu²⁺) having a proportion of 3% Eu and a quantum efficiency of 72%. Figure 7 shows the energy distribution (in arbitrary units) and reflection (in percent) of sample HU 65/00 (Sr₂Si₅N₈:Eu²⁺) having a proportion of 5% Eu and a quantum efficiency of 67%. Figure 8 shows the energy distribution (in arbitrary units) and reflection (in percent) of sample HU 42/00 (Ca₂Si₅N₈:Eu²⁺) having a proportion of 1% Eu and a quantum efficiency of 37%.

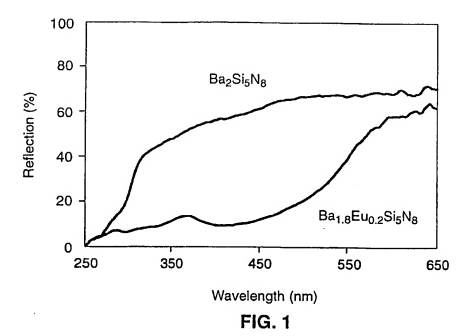
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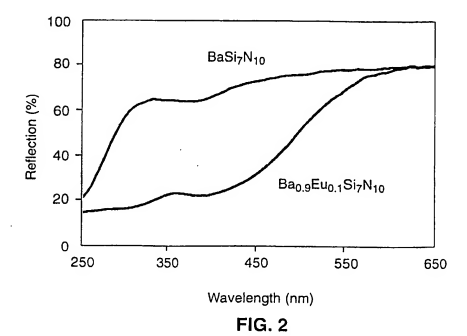
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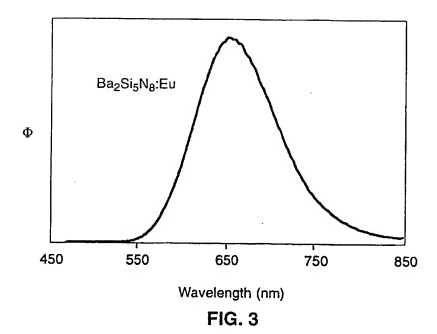
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Claims

- 1. Pigment with day-light fluorescence, characterised in a host lattice of the nitridosilicate type $M_xSi_yN_z$:Eu, wherein M is at least one of an alkaline earth metal chosen from the group Ca, Sr, Ba, Zn and wherein z = 2/3x + 4/3y.
- 2. Pigment according to claim 1, wherein x = 2, and y = 5.
- 5 3. Pigment according to claim 1, wherein x = 1, and y = 7.
 - 4. Pigment according to claim 1, wherein M is strontium.
 - 5. Pigment according to claim 1, wherein M is a mixture of at least two metals of said group.
 - 6. Pigment according to claim 1, wherein Si is replaced fully or partially by Ge.
- 7. Pigment according to claim 1, wherein said pigment is absorbing within the blue to green spectral region.
 - 8. Pigment according to claim 7, wherein said pigment is fluorescent within the yellow to red spectral region.
 - 9. Use of the pigment according to claim 1 as a coloring pigment.
- 15 10. Use of the pigment according to claim 1 as a phosphor excited by light sources, said phosphor emitting within the yellow to red spectral region.

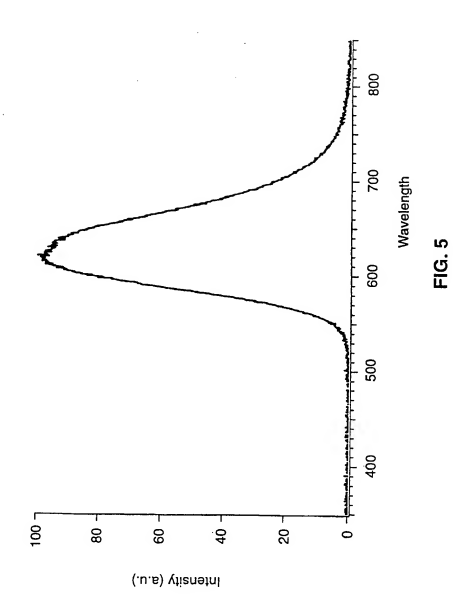


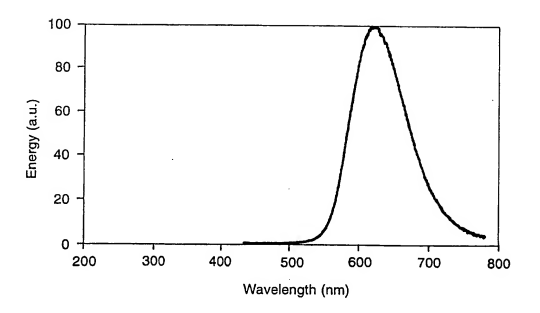




Ф ВаSi₇N₁₀:Eu 450 550 650 750 850 Wavelength (nm)

FIG. 4





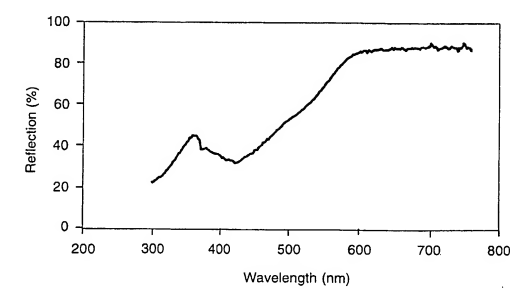
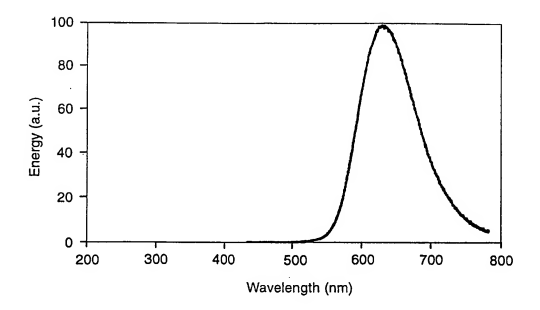


FIG. 6

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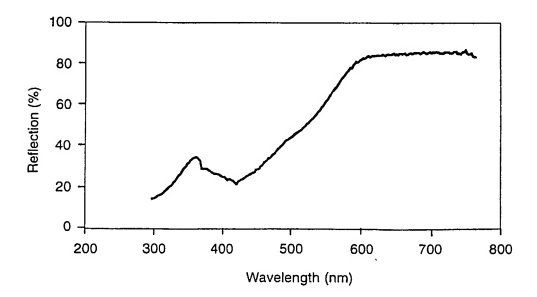
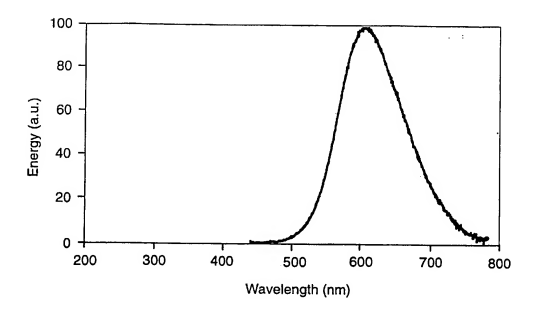


FIG. 7



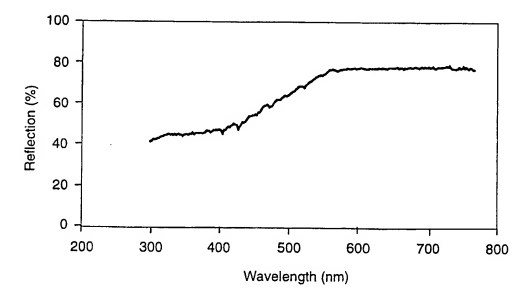


FIG. 8

INTERNATIONAL SEARCH REPORT

In ational Application No PCT/EP 00/12047

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C09K11/79						
According to International Patent Classification (IPC) or to both national classification and IPC						
	SEARCHED					
Minimum do IPC 7	cumentation searched (classification system followed by classificat $C09K$	ion symbols)				
	ion searched other than minimum documentation to the extent that					
Electronic data base consulted during the international search (name of data base and, where practical, search terms used) INSPEC, WPI Data, PAJ, EPO-Internal, IBM-TDB, CHEM ABS Data						
C. DOCUME	ENTS CONSIDERED TO BE RELEVANT					
Category *	Citation of document, with indication, where appropriate, of the re	levant passages	Relevant to claim No.			
A	SOON-SEOK LEE ET AL: "Development luminescent characteristics of Carlo based phosphors" JOURNAL OF THE INSTITUTE OF ELECTENGINEERS OF KOREA D, OCT. 1999, ELECTRON. ENG. KOREA, SOUTH KOREA vol. 36-D, no. 10, pages 31-36, XP002136109 ISSN: 1226-5845 page 31	1-10				
Furti	ner documents are listed in the continuation of box C.	Patent family members are listed	in annex.			
A document defining the general state of the art which is not considered to be of particular relevance *E* earlier document but published on or after the international filing date *L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) *O* document referring to an oral disclosure, use, exhibition or other means *P* document published prior to the international filing date but later than the priority date claimed Date of the actual completion of the international search		"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family Date of mailing of the international search report				
	2 March 2001	19/03/2001 Authorized officer				
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016		Drouot-Onillon, M-C				